

B. Claims

The following is a complete listing of the claims and replaces all earlier listings of claims in the present application.

1. (Currently Amended) A tablet for oral administration that disintegrates quickly in the oral cavity in less than 30 seconds, comprising:

- i) spray-dried mannitol in a proportion of at least 59.5%;
 - ii) active ingredient in a proportion below or equal to 10%, as a fine powder in which at least 90% in weight of the active ingredient has a particle size less than 100 μm ;
 - iii) microcrystalline cellulose in a proportion from 10 to 18%, with an average particle size of approximately 50 μm where at least 99% in weight of microcrystalline cellulose has a particle size below 250 μm ;
 - iv) sodium croscarmellose in a proportion from 1 to 4%; and
 - v) a lubricant agent in a proportion from 0.5 to 2% in weight,
- where, unless specified otherwise, the percentages are expressed in percent weight of the total weight of the tablet, wherein said tablet has a friability below 0.5%.

2. (Cancelled)

3. (Currently Amended) The tablet for oral administration according to claim 1, ~~wherein claim 2, characterised in that~~ it has a friability below 0.2% ~~according to Ph. Eur. 2.9.7.~~

4. (Currently Amended) The tablet for oral administration according to claim 1, ~~characterised in that~~ wherein it has an apparent density from 1.1 to 1.3 g/ml.

5. (Withdrawn - Currently Amended) Tablet for oral administration according to claim 1, ~~characterised in that~~ wherein it has a flavouring agent in a proportion from 0.5 to 2% in weight of the total weight of the tablet.
6. (Withdrawn - Currently Amended) Tablet for oral administration according to claim 5, ~~characterised in that~~ wherein it has an artificial sweetener in a proportion from 0.5 to 2% in weight of the total weight of the tablet.
7. (Withdrawn - Currently Amended) Tablet for oral administration according to claim 1, ~~characterised in that~~ wherein it has a humidity adsorbing agent in a proportion from 0.1 to 0.5% in weight of the total weight of the tablet.
8. (Withdrawn - Currently Amended) Tablet for oral administration according to claim 1, ~~characterised in that~~ wherein it has an anti-adherent agent in a proportion from 0.5 to 2% in weight of the total weight of the tablet.
9. (Currently Amended) The tablet for oral administration according to claim 1, ~~characterised in that the~~ wherein a proportion of insoluble elements is below 20% in weight of the total weight of the tablet.
10. (Withdrawn - Currently Amended) Tablet for oral administration according to ~~any of previous claims, characterised in that it has a round shape, flat, bevelled with~~ claim 1, wherein said tablet has a round shape and is flat and bevelled, said tablet having a thickness from 1.8 to 2.2 mm.

11. (Withdrawn - Currently Amended) Tablet for oral administration according to claim 10, ~~characterised in that~~ wherein it disintegrates quickly in the oral cavity in less than 20 seconds.

12. (Withdrawn - Currently Amended) Process for obtaining a tablet for oral administration as ~~defined in any of claims 1 to 11, characterised in that it comprises~~ comprising the following steps:

- i) ~~Sieving~~ sieving and mixing the components except for the lubricant agent;
- ii) ~~Sieving~~ sieving the lubricant agent;
- iii) ~~Mixing~~ mixing of all the components; and
- iv) ~~Direct compression of~~ directly compressing the final mixture.

13. (Withdrawn - Currently Amended) Process for obtaining a tablet according to claim 12, ~~characterised in that~~ wherein said final mixture has a flowability below or equal to 10 seconds ~~according to Ph. Eur. 2.9.16.~~

14. (Withdrawn - Currently Amended) Process for obtaining a tablet according to claim 12, ~~characterised in that~~ wherein said final mixture has an ability to settle below or equal to 20 ml ~~according to Ph. Eur. 2.9.15.~~